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## PAPER



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A machine learning study on superlattice electron blocking layer design for AlGaN deep ultraviolet light-emitting diodes using the stacked XGBoost/ LightGBM algorithm<sup>†</sup>

Rongyu Lin, <sup>[10]</sup> ‡<sup>a</sup> Zhiyuan Liu, ‡<sup>a</sup> Peng Han, <sup>b</sup> Ronghui Lin, <sup>a</sup> Yi Lu, <sup>a</sup> Haicheng Cao, <sup>a</sup> Xiao Tang, <sup>10</sup> <sup>a</sup> Chuanju Wang, <sup>a</sup> Vishal Khandelwal, <sup>a</sup> Xiangliang Zhang\*<sup>bc</sup> and Xiaohang Li\*<sup>a</sup>

Aluminium gallium nitride (AlGaN)-based deep ultraviolet (DUV) light-emitting diodes (LEDs) suffer from low internal quantum efficiency (IQE) and serious efficiency droop. One reason for this is the electron leakage and poor hole injection related to the band alignment of the heterojunctions, doping, polarization effect, and others. In the past, the AlGaN/AlGaN superlattice (SL) electron blocking layer (EBL) was proposed to optimize the carrier transport and improve the LED performance. However, the SL-EBL design is a trade-off of multiple physics mechanisms, and the LED efficiency deteriorates when the design is improper. We used extreme gradient boosting (XGBoost) and light gradient boosting machine (LightGBM) stacked machine learning (ML) models to predict various high-performance SL-EBLs considering different compositions, thicknesses, and band offset ratios. Based on the ML model, we propose an easier and experimentally achievable low Al-content SL-EBL (1 nm/5 nm  $Al_{0.7}Ga_{0.3}N/$  $Al_{0.58}Ga_{0.42}N$ ) that can significantly optimize carrier transport. The improvement in IQE and wall-plug efficiency could be as high as about 70% compared with those of the conventional bulk EBL. Moreover, we analyze the prediction data and reveal the influence of the composition and thickness on the IQE improvement. The composition difference should be enlarged at a higher band offset ratio, which may be explained by the electron potential and polarization modulation. The critical thickness of the optimized SL-EBL is investigated to guarantee effective electron blocking without destroying the material quality, doping modulation, and operating voltage. This work provides a systematic study of SL-EBLs and helps promote the development and application of SL-EBLs for high-efficiency DUV LEDs.

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## Introduction

Aluminium gallium nitride (AlGaN)-based deep ultraviolet (DUV) light-emitting diodes (LEDs) have undergone rapid development due to their employment in widespread applications, such as water purification, sterilization, communication, and sensing.<sup>1</sup>

Compared with conventional DUV sources, such as mercury lamps, DUV LEDs have obvious advantages in terms of lifetime, size, and environmental protection, and are considered mainstream DUV sources for the future.<sup>2</sup> Although various DUV LED products have been on the market, they usually suffer from low external quantum efficiency.

Transverse magnetic mode light is dominant in DUV emission, which propagates in the lateral direction and is difficult to extract from the device. Thus, the light extraction efficiency of DUV LEDs is usually as low as 10%.<sup>3</sup> In contrast, the nonradiative recombination, polarization effect, poor hole injection, and electron overflow seriously restrict the internal quantum efficiency (IQE) of DUV LEDs.<sup>4–7</sup> High Al-content AlGaN is utilized as the electron blocking layer (EBL) to improve the electron leakage issue. However, the EBL blocks electrons and holes, especially when the valence band offset ( $\Delta E_v$ ) is large.<sup>8</sup> Moreover, the polarization charge at the heterointerface between the last quantum barrier (LQB) and EBL could lead to electron

<sup>&</sup>lt;sup>a</sup> Advanced Semiconductor Laboratory, Computer, Electrical and Mathematical Sciences and Engineering Division, King Abdullah University of Science and Technology (KAUST), Thuwal 23955-6900, Kingdom of Saudi Arabia. E-mail: xiaohang.li@kaust.edu.sa

<sup>&</sup>lt;sup>b</sup> Laboratory Machine, Intelligence and Knowledge Engineering (MINE), Computer, Electrical and Mathematical Sciences and Engineering Division, King Abdullah University of Science and Technology (KAUST), Thuwal 23955-6900, Kingdom of Saudi Arabia

<sup>&</sup>lt;sup>c</sup> Department of Computer Science and Engineering, University of Notre Dame, Notre Dame, IN 46556, USA. E-mail: xzhang33@nd.edu

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<sup>‡</sup> These authors contributed equally.

accumulation.<sup>9</sup> Both shrinking and enlarging the effective electron and hole barrier ( $\Phi_{\rm e}$  and  $\Phi_{\rm h}$ , respectively) compromise the carrier injection efficiency.

Adopting higher p-type doping in the EBL could mitigate this issue and reduce the device resistance simultaneously; however, it is difficult to achieve because of the large activation energy of the Mg acceptor.<sup>10</sup> The energy band can also be modulated by employing various materials and structures. Wen *et al.* proposed an undoped  $B_{0.14}Al_{0.86}N$  EBL to significantly improve electron blocking and hole injection,<sup>11</sup> and Liu *et al.* used InAlN as the LQB to modulate the polarization charge and band bending at the LQB/EBL.<sup>12</sup> Ren *et al.* revealed that the EBL could be removed in DUV LEDs using a composition-grade quantum barrier.<sup>13</sup> Furthermore, the AlN insertion layer, composition-grade LQB and EBL, and N-polar DUV LEDs are demonstrated theoretically and experimentally to improve the efficiency of LEDs.<sup>14–18</sup>

The superlattice (SL) EBL is another promising candidate to replace the bulk EBL. The periodic well/barrier structure provides a multireflection effect, enhancing electron blocking.<sup>19</sup> With the same high barrier, the SL-EBL has a lower average Al content compared with the bulk EBL. Thus, the polarization field and band bending in the LQB could be effectively weakened, contributing to better carrier transport.<sup>20</sup> Numerous reports have found that the short-period SL is helpful for p-type doping, which could further improve the device performance.21-23 Although a graded SL-EBL and an AlInGaN-based SL-EBL are proposed in DUV LEDs for optimization, the conventional AlGaN/AlGaN periodic SL-EBL may be preferred due to the lower growth complexity.<sup>24,25</sup> The design of the SL-EBL is usually dependent on the simulation software. The different combinations of composition and thickness are basic variables contributing to the various device capabilities. However, the polarization charge, potential height, trade-off between electron blocking and hole injection, carrier accumulation, and recombination in the SL must be considered in the SL design.10,26

The physics becomes more complex with the inclusion of the band offset ratio. Moreover, the AlGaN/AlGaN SL-EBL

design principle has scarcely been reported; thus, optimizing the structure perfectly *via* simulations is not straightforward. In this work, we employed stacked extreme gradient boosting (XGBoost) and light gradient boosting machine (LightGBM) machine learning (ML) models to reveal high-performance AlGaN/AlGaN SL-EBL designs in 270 nm DUV LEDs. The model is trained based on the data generated from the software Technology Computer-Aided Design (TCAD), and the accuracy and robustness of the ML model are evaluated using the root mean square error (RMSE). More importantly, with the aid of the ML model, we systematically investigate the dependence of the composition, thickness, and band offset ratio, which could significantly guide the SL-EBL design both theoretically and experimentally in the future.

### Models and parameters

#### Light-emitting diode structures and parameters

As illustrated in Fig. 1(a), the investigated LED architectures are grown on the *c*-plane AlN template. First, a 3 nm-Al<sub>0.6</sub>Ga<sub>0.4</sub>N (Si:  $5 \times 10^{18}$  cm<sup>-3</sup>) is grown as the electron injection layer. This layer is followed by five pairs of AlGaN/AlGaN multi-quantum wells (MQWs) comprising 12 nm Al<sub>0.6</sub>Ga<sub>0.4</sub>N quantum barriers and 3 nm Al<sub>0.5</sub>Ga<sub>0.5</sub>N quantum wells (QWs). Next, different SL-EBLs with 5  $\times$  10<sup>18</sup> cm<sup>-3</sup> p-type doping are simulated for various electron blocking and hole injection capabilities, including a five-pair Alc1Ga1-c1N/Alc2Ga1-c2N SL-EBL. Considering that the EBL should possess a larger bandgap than the OW to block the electrons, the lower limit of the Al content of the barrier  $(c_1)$  and well  $(c_2)$  is set at 50%. The upper limit is 80%, depending on the growth and doping difficulty of AlGaN with a high Al content.<sup>27,28</sup> The thickness of the barrier  $(t_1)$  and well  $(t_2)$  vary from 1 to 5 nm. Finally, 100 nm p-Al<sub>0.6</sub>Ga<sub>0.4</sub>N (Mg:  $3 \times 10^{19} \text{ cm}^{-3}$ ) and 50 nm p-GaN (Mg:  $1 \times 10^{20} \text{ cm}^{-3}$ ) are deposited as the hole injection layer and cap layer, respectively. The mesa size of the devices in the simulation is 300  $\mu$ m imes300 µm, and the expected wavelength is 270 nm.

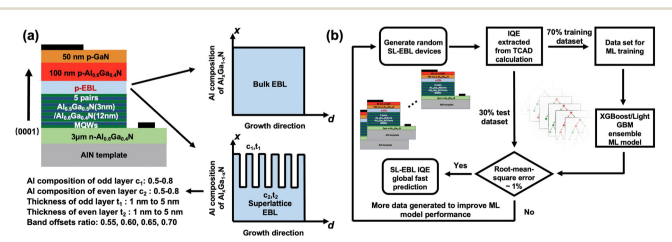


Fig. 1 (a) Cross-sectional diagram of light-emitting diode (LED) structures. (b) Workflow chart of the extreme gradient boosting (XGBoost) and light gradient boosting machine (LightGBM) stacked machine learning (ML) model prediction method.

In this study, the APSYS TCAD software by Crosslight is employed to consistently solve various physical equations and provide adequate data for ML.<sup>29</sup> The operating temperature and background loss are set at 300 K and 2000 m<sup>-1</sup>, respectively.<sup>30,31</sup> The auger recombination coefficient, Shockley-Read-Hall recombination lifetime, and radiative recombination coefficient are estimated respectively to be  $2.88 \times 10^{-30}$  cm<sup>6</sup> s<sup>-1</sup>, 15 ns, and  $2.13 \times 10^{-11}$  cm<sup>3</sup> s<sup>-1</sup>.<sup>32</sup> The percentage of the polarization charge screened by the defects and carrier injection is set at 50%.33 The band offset ratios of AlGaN/AlGaN heterojunctions rely on material growth conditions and influence the simulation result. According to previous reports, diverse values from 0.55:0.45 to 0.70:0.30 are considered in this work to offer a broad reference for the SL design.<sup>34</sup> Other parameters are set as the default values in APSYS, so the results are reproducible by others.

# Machine learning method for the superlattice electron blocking layer design

**Data preparation.** Fig. 1(b) illustrates the workflow of the ML model construction. The ultraviolet (UV) LEDs with the various mentioned EBL designs are calculated using TCAD software. To build an efficient ML model, we split the TCAD calculation results into 70% for the training dataset and 30% for the testing dataset. The TCAD prediction requires substantial calculation resources; therefore, we used the try-and-catch method, where we first constructed the ML model with a small volume of TCAD calculation data and added more data from new calculations to enhance the ML model until the performance reached a specific criterion. The details of the parameters for the ML model are listed in Table 1.

XGBoost/LightGBM stacking ensemble machine learning model. ML is increasingly employed in semiconductor materials and devices for mechanism investigation and structure design. Recently, researchers used a neural network structure to predict the optical and electronic properties of semiconductor devices.<sup>35–39</sup> Lin *et al.* applied a tree-based ensemble XGBoost model to predict the tunnel junction resistance and designed a novel asymmetrical tunnel junction with low resistance.<sup>40</sup> In this paper, we propose a new ML framework for the SL-EBL design, which can predict the IQE with high accuracy. Instead of incurring a single model, we stacked two different existing methods in the training and testing process: XGBoost and LightGBM.<sup>41,42</sup> In this framework, we use *m* XGBoost models and *n* LightGBM models and obtain results by stacking them. Given the *i*th sample's feature  $x_i$ , we use  $f_j^g(x_i)$  to denote the result of the *j*th XGBoost model with the corresponding

Table 1Parameters of the extreme gradient boosting (XGBoost)/lightgradient boosting machine (LightGBM) stacked machine learning (ML)model

	Number of training data		т	n	α	β
0.55	691	298	1	2	0.4359	0.4872/0.0769
0.6	416	180	2	1	0.1905/0.2381	0.5714
0.65	700	301	1	2	0.4536	0.5361/0.0103
0.7	465	200	2	1	0.4714/0.0143	0.5143

weight  $\alpha_{j}$ , and  $f_{j}^{l}(x_{i})$  to denote the result of the *j*th LightGBM model with the corresponding weight  $\beta_{j}$ . Then, the final predicted result  $\hat{y}_{i}$  of the *i*th sample is given as follows:

$$\hat{y}_i = \sum_{j=1}^m \alpha_j f_j^{g}(x_i) + \sum_{j=1}^n \beta_j f_j^{1}(x_i)$$
(1)

The objective functions of XGBoost and LightGBM are the same. Both are inherited from the gradient boosting decision tree method, which is optimized iteratively, with the addition of a regularization term. Taking the XGBoost method as the example, given the corresponding features  $\{x_1, x_2, ..., x_n\}$  and prediction results  $\{y_1, y_2, ..., y_n\}$ , the objective function  $Obj^{(1)}$  of the first iteration in XGBoost is as follows:

$$Obj^{(1)} = \sum_{i=1}^{n} l\left(y_i, f^{g}_{(1)}(x_i)\right) + \Omega\left(f^{g}_{(1)}\right)$$
(2)

where  $l(y_i, f_{(1)}^g(x_i)) = ||y_i - f_{(1)}^g(x_i)||_2^2$  is used to compute the difference between  $y_i$  and  $f_{(1)}^g(x_i)$ , and  $\Omega(f_{(1)}^g)(x_i)$  represents the regularization term constraining the parameters in the regression function  $f_{(1)}^g$ . Then, for the *t*th iteration, we obtain the objective function Obj<sup>(t)</sup> of XGBoost as follows:

$$Obj^{(t)} = \sum_{i=1}^{n} l\left(y_i, \sum_{j=1}^{t-1} f_{(j)}^{g}(x_i) + f_{(t)}^{g}(x_i)\right) + \Omega\left(f_{(t)}^{g}\right)$$
(3)

Then, after *T* iterations, we obtain the predicted IQE of the *i*th sample for the XGBoost method as  $f^{g}(x_i) = \sum_{j=1}^{T} f^{g}_{(j)}(x_i)$ . The objective function for the LightGBM is the same as for XGBoost, which uses different optimization methods from XGBoost in the training process.

#### Evaluation of the proposed machine learning model

The RMSE values are calculated to illustrate the performance of the proposed ML model, which is relatively intuitive to the magnitude of IQE. The mathematical expression of RMSE is given as follows:

$$\mathbf{RMSE} = \sqrt{\frac{\sum\limits_{i=1}^{N} \left( \mathrm{IQE}_{\mathrm{ML \ prediction}} - \mathrm{IQE}_{\mathrm{TCAD \ prediction}} \right)^{2}}{N}} \quad (4)$$

where *N* is the number of IQE data in the testing dataset. With this framework, we can combine XGBoost and LightGBM to predict the IQE, and their influence is controlled by the hyperparameters *m*, *n*,  $\alpha$ , and  $\beta$ . Combining these two models makes this framework fit the training data better than the single model. In Fig. 2, IQEs from the ML prediction are compared to the IQE from the testing dataset, and the scatter points are well located along the line *y* = *x*, demonstrating the feasibility of the ML model. The regularization term in the objective function can overcome the overfitting problem well. The proposed models exhibit low RMSEs, which are 1.6%, 1.7%, 1.1%, and 0.9% for the band offsets of 0.55, 0.6, 0.65, and 0.7, respectively.

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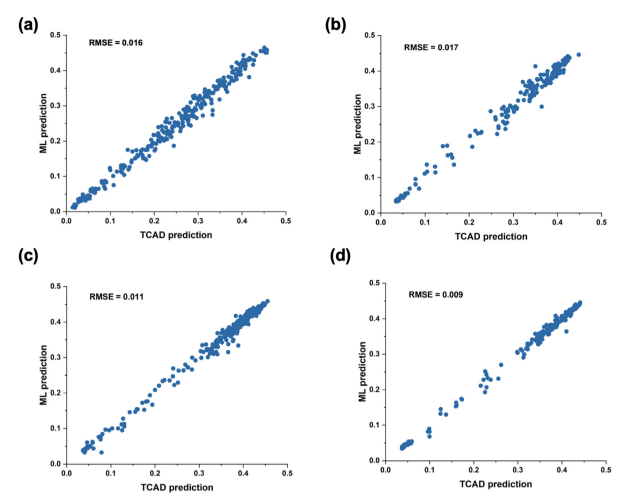


Fig. 2 Internal quantum efficiency values from the machine learning (ML) prediction *versus* the TCAD prediction of the band offsets = (a) 0.55 (b) 0.6 (c) 0.65 and (d) 0.7.

#### Results and discussion

We plotted IQEs for changing  $c_1$  and  $c_2$  at an odd and even number of layer thicknesses and band offsets to better understand the results. The values for  $t_1$  and  $t_2$  increase from 1 to 5 nm with a step of 1 nm, and the band offsets vary from 0.55 to 0.7 with 0.05 intervals. The details for all figures are listed in the ESI.†

The ESI<sup>†</sup> provides Al compositions, thicknesses of odd and even-numbered layers, and the band offsets, which affect the performance of UV LEDs. Thus, a detailed investigation is conducted to understand the correlation between SL-EBL designs and UV LED performance.

# Low Al-content superlattice electron blocking layer design from machine learning prediction

High p-type doping can be easily achieved with low Al content AlGaN because of the lower Mg activation energy, and the higher Al incorporation efficiency can facilitate epitaxy growth.<sup>43,44</sup> Thus, predicting high-efficiency DUV LEDs with relatively low Al content, such as lower than 70%, is meaningful for practical devices. We selected a low Al-content SL-EBL (Sample A) from the predictions, which is compared to Al<sub>0.7</sub>Ga<sub>0.3</sub>N (Sample B) and  $Al_{0.8}Ga_{0.2}N$  (Sample C) bulk EBLs with the same total thickness. The compositions of the barrier and well are 0.70 and 0.58, and the thickness is 1 and 5 nm, respectively. The discussed band offset ratio is 0.6. All designs at different band offset ratios are presented in the ESI<sup>†</sup> and are not listed here.

Fig. 3(a) illustrates that Sample A has the highest IQE compared with Samples B and C, and the efficiency droop is also significantly suppressed. By appropriately designing the SL-EBL, DUV LEDs could perform better with less epitaxy difficulty. Using Fig. 3(b), we analyzed the electron and hole density lateral distribution at the interface between the EBL and MQWs, reflecting the electron leakage from MQWs to p-layers and hole injection from p-layers to MQWs. The figure reveals that Sample A has the lowest electron current, whereas Samples B and C face more serious electron leakage. Although Sample C has the highest hole current peak for the hole injection, Sample A has a more uniform hole current distribution along the mesa. Thus, the IQE improvement of the SL-EBL DUV LED can be concluded due to the better electron blocking and hole injection capability. Fig. 4 and Table 2 present the I-V curve and wall-plug efficiency (WPE) of Samples A, B, and C. Among the three structures, Sample C has the largest operating voltage due to the higher

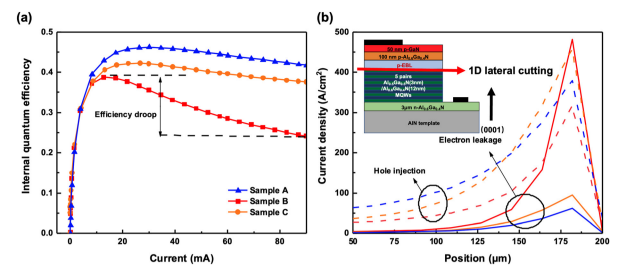


Fig. 3 (a) Internal quantum efficiency curve of deep ultraviolet light-emitting diodes with bulk electron blocking layers (EBLs) and superlattice EBLs. (b) Electron and hole (dash line) current density *versus* lateral position at the interface between the EBL and MQWs.

electron and hole potential from the high Al-content EBL, which compromises the WPE.

The situation of Samples A and B is more complex. In the beginning, Sample A has a lower operating voltage (open voltage) than Sample B because the SL-EBL has a lower average Al content, leading to a lower potential difference. At the higher current, the operating voltage is higher in Sample A (5.52 V at 90 mA) compared with Sample B (5.38 V at 90 mA). The periodic barrier well structure of SL-EBL could confine carriers and block the carrier transport, corresponding to a higher voltage at the same current. Finally, Sample A presents the highest WPE (3.48% at 90 mA) because of the excellent IQE and medium operating voltage.

Composition design under different band offset ratios

From the data in the ESI,<sup>†</sup> it can be observed that the IOE

highly depends on the composition of the SL-EBL. The highest

#### Table 2 Operating voltage and wall-plug efficiency at 90 mA

	Operating voltage (V)	Wall-plug efficiency (%)
Al <sub>0.7</sub> Ga <sub>0.3</sub> N bulk EBL (Sample B)	5.38	2.06
Al <sub>0.8</sub> Ga <sub>0.2</sub> N bulk EBL (Sample C)	5.70	3.03
Al <sub>0.7</sub> Ga <sub>0.3</sub> N/Al <sub>0.58</sub> Ga <sub>0.42</sub> N SL-EBL (Sample A)	5.52	3.48

Notes: EBL: electron blocking layer; SL: superlattice

IQE always occurs in the region where the SL-EBL possesses the  $Al_{0.8}Ga_{0.2}N$  SL barrier. The outcome is easily understood because the higher Al-content barrier can provide a stronger electron barrier to suppress electron leakage, and the polarization charge and band can be modulated by employing the SL well with different compositions. We investigate the influence

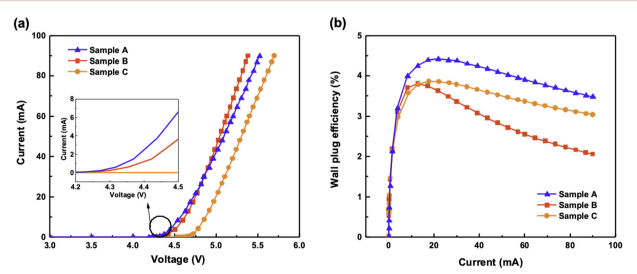


Fig. 4 (a) Current–voltage (*I–V*) curve and (b) wall-plug efficiency of deep ultraviolet light-emitting diodes with bulk electron blocking layers (EBLs) and superlattice EBLs.

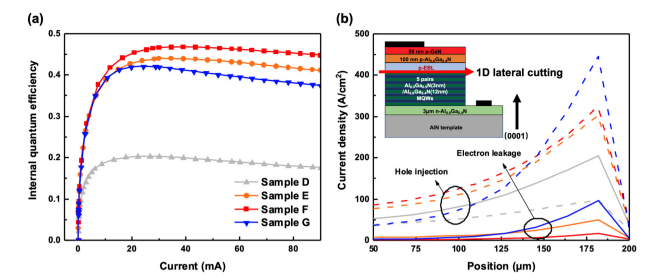


Fig. 5 (a) Internal quantum efficiency curve of samples. (b) Electron and hole current density *versus* lateral position at the interface between EBL and MQWs.

of the SL composition on the IQE. The barrier composition is locked as 0.8 with well compositions of 0.5 (Sample D), 0.6 (Sample E), 0.7 (Sample F), and 0.8 (Sample G). The discussed thickness is  $t_1 = t_2 = 3$  nm, and the band offset ratio is set to 0.6.

Fig. 5 depicts the calculated IQE curve for different samples. Sample F has the highest IQE, whereas Sample D has the lowest IQE and worse performance than the bulk EBL LED. The electron and hole lateral density distribution along the interface between MQWs and EBL is shown in Fig. 5(b). Sample F has the lowest electron leakage of all samples, and the hole injection is strong and uniform. However, Sample D suffers from poor carrier transport. Fig. 6 presents the radiative recombination rate distribution of Samples D and F to further illustrate the carrier transport deficiency in Sample D. In addition to the recombination in the MQWs, Sample D displays a strong recombination in the SL well. Due to the same composition as QWs, the SL wells play the role of extra QWs in UV LEDs. They are sandwiched between a higher Al-content barrier (Al<sub>0.8</sub>Ga<sub>0.2</sub>N), with stronger confinement for carriers. Thus, electrons in the MQWs prefer transferring to the SL and recombining with holes in the SL well. The recombination in the MQWs of Sample D is much weaker than that in Sample F, resulting in poor IQE.

As mentioned, the band offset ratio of AlGaN significantly influences the carrier transport. Although the band offset ratio of AlGaN/AlGaN heterojunctions is normally considered as a fixed number under specific growth conditions, it can be modulated by the interfacial structure, defect position, interface states and Al content, which result from different epitaxy conditions.<sup>45</sup> Thus, it is necessary to understand the influence of the band offset ratio on the SL-EBL design. In other words, the same design could not be directly transferred under dissimilar growth conditions. In Fig. 7, we set  $c_1$  to 0.8 and predict

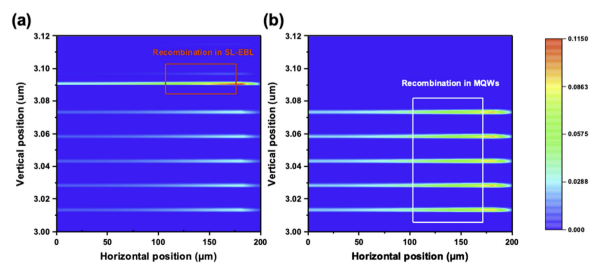


Fig. 6 Radiative recombination rate for (a) Sample D and (b) Sample F.

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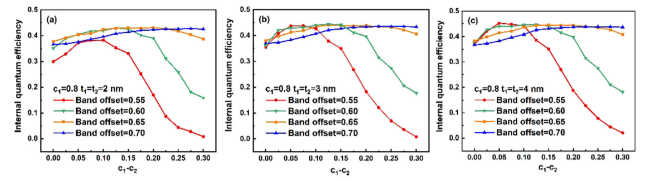


Fig. 7 Internal quantum efficiency (IQE) at 90 mA with various composition difference  $(c_1 - c_2)$  superlattice electron blocking layers at band offset ratios of  $t_1 = t_2$  at (a) 2 nm, (b) 3 nm, (c) and 4 nm.

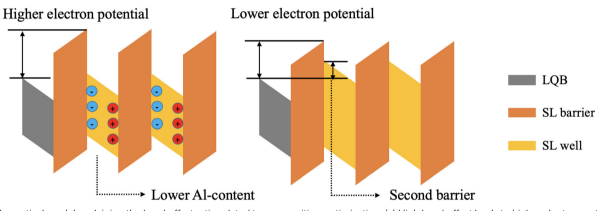


Fig. 8 Theoretical model explaining the band offset ratio related to composition optimization. (a) High band offset leads to higher electron potential. (b) Low band offset leads to low electron potential between the multi-quantum well and superlattice electron blocking layer.

the IQE with various well-barrier composition differences. Furthermore, various band offset ratios and thicknesses are considered, revealing that the overall trends are similar regardless of the thickness of SL-EBLs. The IQE peak occurs at a larger composition difference when the band offset ratio increases, also shown in the ESI,† where generally the red region gradually shifts from the upper right corner to the upper left and lower right directions in the figures, indicating that the lower Al

content of the SL well is desirable for improving the IQE. To explain this result, we use a trade-off model between polarization modulation and electron potential. As illustrated in Fig. 8, the electron potential is also lower at the lower band offset ratio due to the limited conduction band offset. Hence, the SL well should maintain a higher Al content to guarantee sufficient electron blocking capability. The electron potential is high enough for the larger band offset ratio, so it is better to use a low Al-content SL

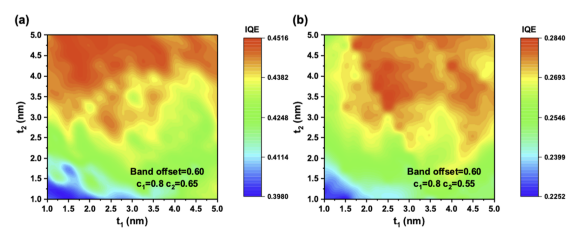


Fig. 9 (a) Thickness to internal quantum efficiency relationship for (a)  $c_1 = 0.8$  and  $c_2 = 0.55$  and (b)  $c_1 = 0.8$  and  $c_2 = 0.65$ .

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well to reduce the average Al content and modulate the polarization charge. We believe that this conclusion can significantly guide SL-EBL design.

#### Thickness factors on superlattice electron blocking layer design

Fig. 9 presents the thickness influence on the IQE of the SL-EBL DUV LEDs under two well compositions. The optimal thickness simultaneously depends on the SL composition and band offset ratio. However, the IQE is compromised if the well and barrier are extremely thin, such as 1 and 2 nm, because of the poor electron blocking ability. In contrast, compared with the medium thickness area, a too-thick SL, such as 5 nm, has no obvious advantage in IQE.

Moreover, a too-thick SL also leads to various problems. First, a too-thick SL can deteriorate the material quality because of the lattice mismatch. Second, p-type doping modulation is stronger in a short-period SL. Third, the operating voltage increases with the SL thickness, shirking the WPE and accumulating more heat. Thus, the first consideration of the design should focus on a medium region, such as a 3 nm average thickness.

## Conclusions

In summary, to the best of our knowledge, a novel stacked XGBoost/LightGBM ML model is used to efficiently predict the UV LED IQE performance. The ML model is evaluated by its RMSEs and the low error values (around  $\sim 1\%$ ), demonstrating that it is promising and accurate in terms of IQE predictions. Then, a systematic investigation of SL-EBL is conducted based on the ML model predictions, and a high-efficiency design can be achieved through the results. The improvement of IQE originates from the better electron blocking and hole injection, which were analyzed using the horizontal current distribution and radiative recombination rate. The optimized low Al-content SL-EBL is proposed to reduce epitaxy complexity, and it induces a nearly 70% IQE and WPE improvement compared with the Al<sub>0.7</sub>Ga<sub>0.3</sub>N bulk EBL. Furthermore, the physics mechanism of the SL-EBL was also systemically studied, including the influence of the composition, thickness, and band offset ratio. The composition difference between the SL well and barrier should be increased at larger band offset ratios, which can be explained by the trade-off between the electron potential and polarization modulation. The SL thickness is critical, and an average thickness of about 3 to 4 nm is recommended. We believe that the proposed ML method can also be applied in other LED structure investigations, and this study could deepen the understanding of the SL-EBL and guide the design theoretically and experimentally, boosting the development of high-efficiency DUV LEDs in the future.

## Conflicts of interest

There are no conflicts to declare.

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